

# Package ‘nftbart’

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**Type** Package

**Title** Nonparametric Failure Time Bayesian Additive Regression Trees

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**Description** Nonparametric Failure Time (NFT) Bayesian Additive Regression Trees (BART): Time-to-event Machine Learning with Heteroskedastic Bayesian Additive Regression Trees (HBART) and Low Information Omnibus (LIO) Dirichlet Process Mixtures (DPM). An NFT BART model is of the form  $Y = \mu + f(x) + sd(x) E$  where functions  $f$  and  $sd$  have BART and HBART priors, respectively, while  $E$  is a nonparametric error distribution due to a DPM LIO prior hierarchy. See the following for a complete description of the model at <doi:10.1111/biom.13857>.

**License** GPL (>= 2)

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bartModelMatrix	<i>Deprecated: use bMM instead</i>
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## Description

Create a matrix out of a vector or data.frame. The compiled functions of this package operate on matrices in memory. Therefore, if the user submits a vector or data.frame, then this function converts it to a matrix. Also, it determines the number of cutpoints necessary for each column when asked to do so.

## Usage

```
bartModelMatrix(X, numcut=0L, usequants=FALSE, type=7, rm.const=FALSE,
                cont=FALSE, xicuts=NULL, rm.vars=NULL)
```

## Arguments

X	A vector or data.frame to create the matrix from.
numcut	The maximum number of cutpoints to consider. If numcut=0, then just return a matrix; otherwise, return a list.
usequants	If usequants is FALSE, then the cutpoints in xinfo are generated uniformly; otherwise, if TRUE, quantiles are used for the cutpoints.
type	Determines which quantile algorithm is employed.
rm.const	Whether or not to remove constant variables.
cont	Whether or not to assume all variables are continuous.
xicuts	To specify your own cut-points, use the xicuts argument.
rm.vars	The variables that you want removed.

## Value

If numcut==0 (the default), then a matrix of the covariates is returned; otherwise, a list is returned with the following values.

X	A matrix of the covariates with n rows and p columns.
numcut	A vector of length p of the number of cut-points for each covariate.

`grp` A vector that corresponds to variables in the input `data.frame` that were translated into dummy columns in the output matrix, i.e., for each input variable in order, there is a number in the vector corresponding to the number of output columns created for it.

### See Also

[bMM](#)

### Examples

```
## set.seed(99)

## a <- rbinom(10, 4, 0.4)

## table(a)

## x <- runif(10)

## df <- data.frame(a=factor(a), x=x)

## (b <- bartModelMatrix(df))

## (b <- bartModelMatrix(df, numcut=9))

## (b <- bartModelMatrix(df, numcut=9, usequants=TRUE))

## Not run:
## this is an error
## f <- bartModelMatrix(as.character(a))

## End(Not run)
```

---

bMM

*Create a matrix out of a vector or data.frame*

---

### Description

Adapted from `bartModelMatrix()`. The compiled functions of this package operate on matrices in memory. Therefore, if the user submits a vector or `data.frame`, then this function converts it to a matrix. Also, it determines the number of cutpoints necessary for each column when asked to do so.

### Usage

```
bMM(X, numcut=0L, usequants=FALSE, type=7, xicuts=NULL, rm.const=FALSE,
     rm.dupe=FALSE, method="spearman", use="pairwise.complete.obs")
```

**Arguments**

<code>X</code>	A vector or data.frame to create the matrix from.
<code>numcut</code>	The maximum number of cutpoints to consider. If <code>numcut=0</code> , then just return a matrix; otherwise, return a list.
<code>usequants</code>	If <code>usequants</code> is <code>FALSE</code> , then the cutpoints in <code>xinfo</code> are generated uniformly; otherwise, if <code>TRUE</code> , quantiles are used for the cutpoints.
<code>type</code>	Determines which quantile algorithm is employed.
<code>xicuts</code>	To specify your own cut-points, use the <code>xicuts</code> argument.
<code>rm.const</code>	To remove constant variables or not.
<code>rm.dupe</code>	To remove duplicate variables or not.
<code>method, use</code>	Correlation options.

**Value**

If `numcut==0` (the default), then a matrix of the covariates is returned; otherwise, a list is returned with the following values.

<code>X</code>	A matrix of the covariates with <code>n</code> rows and <code>p</code> columns.
<code>numcut</code>	A vector of length <code>p</code> of the number of cut-points for each covariate.
<code>grp</code>	A vector that corresponds to variables in the input data.frame that were translated into dummy columns in the output matrix, i.e., for each input variable in order, there is a number in the vector corresponding to the number of output columns created for it.
<code>dummy</code>	Corresponds to <code>grp</code> with a two row matrix including the start column of each dummy group in row 1 and the end column in row 2.

**See Also**

[xicuts](#)

**Examples**

```
set.seed(99)

a <- rbinom(10, 4, 0.4)

table(a)

x <- runif(10)

df <- data.frame(a=factor(a), x=x)

(b <- bMM(df))

(b <- bMM(df, numcut=9))
```

```
(b <- bMM(df, numcut=9, usequants=TRUE))  
  
## Not run:  
  ## this is an error  
  f <- bMM(as.character(a))  
  
## End(Not run)
```

---

bmx

*NHANES 1999-2000 Body Measures and Demographics*

---

### Description

This data set was created from the National Health and Nutrition Examination Survey (NHANES) 1999-2000 Body Measures Exam and Demographics. To create growth charts, this data is restricted to 3435 children aged 2 to 17.

### Usage

```
data(bmx)
```

### Format

SEQN:	Sequence number
BMXHT:	Height in cm
RIAGENDR:	Gender: 1=male, 2=female
RIDAGEEX:	Age in years with fractions for months
RIDRETH2:	Race/ethnicity: 1=Non-Hispanic White, 2=Non-Hispanic Black, 3=Hispanic
BMXWT:	Weight in kg

### References

National Health and Nutrition Examination Survey 1999-2000 Body Measures Exam. <https://www.cdc.gov/Nchs/Nhanes/1999-2000/BMX.htm>

National Health and Nutrition Examination Survey 1999-2000 Demographics. <https://www.cdc.gov/Nchs/Nhanes/1999-2000/DEMO.htm>

---

CDHeight

*CDC height for age growth charts*

---

**Description**

Using the Cole and Green LMS method, here we provide percentiles of height by age and sex based on the US National Center for Health Statistics data for children aged 2 to 17.

**Usage**

```
data(CDCheight)
```

**Format**

age:	Age in years
sex:	1=male, 2=female
height.XXX:	Height XXXth percentile in cm

**References**

Cole, Timothy J and Green, Pamela J (1992) Smoothing reference centile curves: the LMS method and penalized likelihood. *Statistics in medicine*, **11**, 1305–1319.

The US Centers for Disease Control and Prevention stature by age LMS parameters <https://www.cdc.gov/growthcharts/data/zscore/statage.csv>

---

CDimpute

*Cold-deck missing imputation*

---

**Description**

This function imputes missing data.

**Usage**

```
CDimpute(x.train, x.test=matrix(0, 0, 0), impute.bin=NULL)
```

**Arguments**

x.train	The training matrix.
x.test	The testing matrix, if given.
impute.bin	An index of the columns to avoid imputing which will be handled by BART internally.

**Details**

We call this method cold-decking in analogy to hot-decking. Hot-decking was a method commonly employed with US Census data in the early computing era. For a particular respondent, missing data was imputed by randomly selecting from the responses of their neighbors since it is assumed that the values are likely similar. In our case, we make no assumptions about which values may, or may not, be nearby. We simply take a random sample from the matrix rows to impute the missing data. If the training and testing matrices are the same, then they receive the same imputation.

**Value**

<code>x.train</code>	The imputed training matrix.
<code>x.test</code>	The imputed testing matrix.
<code>miss.train</code>	A summary of the missing variables for training.
<code>miss.test</code>	A summary of the missing variables for testing.
<code>impute.flag</code>	Whether <code>impute.bin</code> columns were, or were not, imputed.
<code>same</code>	Whether <code>x.train</code> and <code>x.test</code> are identical.

---

Cindex

---

*Calculate the C-index/concordance for survival analysis.*


---

**Description**

The C-index for survival analysis is the corollary of the c statistic (the area under the Receiver Operating Characteristic curve) for binary outcomes. As a probability, the higher is the C-index, the better is the model discrimination vs. lesser probability values. Similarly, the concordance is calculated like the C-index from z-draws via the posterior predictive distribution restricted to the horizon of the data (a la restricted mean survival time).

**Usage**

```
Cindex(risk, times, delta=NULL)
```

```
concordance(draws, times, delta=NULL)
```

**Arguments**

<code>risk</code>	A vector or prognostic risk scores.
<code>draws</code>	A vector of draws via the posterior predictive distribution restricted to the horizon of the data (a la restricted mean survival time).
<code>times</code>	A vector of failure times.
<code>delta</code>	The corresponding failure time status code: 0, right-censored; 1, failure; or 2, left-censored. Defaults to all failures if not specified.

**Value**

The return value is the calculated C-index/concordance.

**References**

Harrell FE, Califf RM, Pryor DB, Lee KL, Rosati RA. (1982) Evaluating the yield of medical tests. JAMA, May 14;247(18):2543-6.

**See Also**

[predict.nft](#)

**Examples**

```
data(lung)
N=length(lung$status)

##lung$status: 1=censored, 2=dead
##delta: 0=censored, 1=dead
delta=lung$status-1

## this study reports time in days
times=lung$time
times=times/7 ## weeks

## matrix of covariates
x.train=cbind(lung[, -(1:3)])
## lung$sex:      Male=1 Female=2

## Not run:
set.seed(99)
post=nft(x.train, times, delta, K=0)
pred=predict(post, x.train, XPtr=TRUE, seed=21)
print(Cindex(pred$logt.test.mean, times, delta))

## End(Not run)
```

---

lung

*NCCTG Lung Cancer Data*

---

**Description**

Survival for 228 patients with advanced lung cancer was recorded up to a median of roughly one year by the North Central Cancer Treatment Group. Performance scores rate how well the patient can perform usual daily activities.

**Format**



inst: Institution code  
 time: Survival time in days  
 status: censoring status 1=censored, 2=dead  
 age: Age in years  
 sex: Male=1 Female=2  
 ph.ecog: ECOG performance score (0=good 5=dead)  
 ph.karno: Karnofsky performance score (bad=0-good=100) rated by physician  
 pat.karno: Karnofsky performance score as rated by patient  
 meal.cal: Calories consumed at meals  
 wt.loss: Weight loss in last six months

### Source

Terry Therneau

### References

Loprinzi CL. Laurie JA. Wieand HS. Krook JE. Novotny PJ. Kugler JW. Bartel J. Law M. Bateman M. Klatt NE. et al. Prospective evaluation of prognostic variables from patient-completed questionnaires. North Central Cancer Treatment Group. *Journal of Clinical Oncology*. 12(3):601-7, 1994.

### Examples

```
data(lung)
```

---

nft2

*Fit NFT BART models.*

---

### Description

The `nft2()/nft()` function is for fitting NFT BART (Nonparametric Failure Time Bayesian Additive Regression Tree) models with different train/test matrices for  $f$  and  $sd$  functions.

### Usage

```

nft2(
  ## data
  xftrain, xstrain, times, delta=NULL,
  xftest=matrix(nrow=0, ncol=0),
  xstest=matrix(nrow=0, ncol=0),
  rm.const=TRUE, rm.dupe=TRUE,
  ## multi-threading
  tc=getOption("mc.cores", 1),

```

```

##MCMC
nskip=1000, ndpost=2000, nadapt=1000, adaptevery=100,
chvf=NULL, chvs=NULL,
method="spearman", use="pairwise.complete.obs",
pbd=c(0.7, 0.7), pb=c(0.5, 0.5),
stepwpert=c(0.1, 0.1), probchv=c(0.1, 0.1),
minnumbot=c(5, 5),
## BART and HBART prior parameters
ntree=c(50, 10), numcut=100,
xifcuts=NULL, xiscuts=NULL,
power=c(2, 2), base=c(0.95, 0.95),
## f function
fmu=NA, k=5, tau=NA, dist='weibull',
## s function
total.lambda=NA, total.nu=10, mask=NULL,
## survival analysis
K=100, events=NULL, TSVS=FALSE,
## DPM LIO
drawDPM=1L,
alpha=1, alpha.a=1, alpha.b=0.1, alpha.draw=1,
neal.m=2, constrain=1,
m0=0, k0.a=1.5, k0.b=7.5, k0=1, k0.draw=1,
a0=3, b0.a=2, b0.b=1, b0=1, b0.draw=1,
## misc
na.rm=FALSE, probs=c(0.025, 0.975), printevery=100,
transposed=FALSE, pred=FALSE
)

nft(
## data
x.train, times, delta=NULL, x.test=matrix(nrow=0, ncol=0),
rm.const=TRUE, rm.dupe=TRUE,
## multi-threading
tc=getOption("mc.cores", 1),
##MCMC
nskip=1000, ndpost=2000, nadapt=1000, adaptevery=100,
chv=NULL,
method="spearman", use="pairwise.complete.obs",
pbd=c(0.7, 0.7), pb=c(0.5, 0.5),
stepwpert=c(0.1, 0.1), probchv=c(0.1, 0.1),
minnumbot=c(5, 5),
## BART and HBART prior parameters
ntree=c(50, 10), numcut=100, xicuts=NULL,
power=c(2, 2), base=c(0.95, 0.95),
## f function
fmu=NA, k=5, tau=NA, dist='weibull',
## s function
total.lambda=NA, total.nu=10, mask=NULL,

```

```

## survival analysis
K=100, events=NULL, TSVS=FALSE,
## DPM LIO
drawDPM=1L,
alpha=1, alpha.a=1, alpha.b=0.1, alpha.draw=1,
neal.m=2, constrain=1,
m0=0, k0.a=1.5, k0.b=7.5, k0=1, k0.draw=1,
a0=3, b0.a=2, b0.b=1, b0=1, b0.draw=1,
## misc
na.rm=FALSE, probs=c(0.025, 0.975), printevery=100,
transposed=FALSE, pred=FALSE
)

```

### Arguments

<code>xftrain</code>	<code>n x pf</code> matrix of predictor variables for the training data.
<code>xstrain</code>	<code>n x ps</code> matrix of predictor variables for the training data.
<code>x.train</code>	<code>n x p</code> matrix of predictor variables for the training data.
<code>times</code>	<code>n x 1</code> vector of the observed times for the training data.
<code>delta</code>	<code>n x 1</code> vector of the time type for the training data: 0, for right-censoring; 1, for an event; and, 2, for left-censoring.
<code>xfctest</code>	<code>m x pf</code> matrix of predictor variables for the test set.
<code>xstest</code>	<code>m x ps</code> matrix of predictor variables for the test set.
<code>x.test</code>	<code>m x p</code> matrix of predictor variables for the test set.
<code>rm.const</code>	To remove constant variables or not.
<code>rm.dupe</code>	To remove duplicate variables or not.
<code>tc</code>	Number of OpenMP threads to use.
<code>nskip</code>	Number of MCMC iterations to burn-in and discard.
<code>ndpost</code>	Number of MCMC iterations kept after burn-in.
<code>nadapt</code>	Number of MCMC iterations for adaptation prior to burn-in.
<code>adaptevery</code>	Adapt MCMC proposal distributions every <code>adaptevery</code> iteration.
<code>chvf, chvs, chv</code>	Predictor correlation matrix used as a pre-conditioner for MCMC change-of-variable proposals.
<code>method, use</code>	Correlation options for change-of-variable proposal pre-conditioner.
<code>pbd</code>	Probability of performing a birth/death proposal, otherwise perform a rotate proposal.
<code>pb</code>	Probability of performing a birth proposal given that we choose to perform a birth/death proposal.
<code>stepwpert</code>	Initial width of proposal distribution for perturbing cut-points.
<code>probchv</code>	Probability of performing a change-of-variable proposal. Otherwise, only do a perturb proposal.

minnumbot	Minimum number of observations required in leaf (terminal) nodes.
nrtree	Vector of length two for the number of trees used for the mean model and the number of trees used for the variance model.
numcut	Number of cutpoints to use for each predictor variable.
xifcuts,xiscuts,xicuts	More detailed construction of cut-points can be specified by the xicuts function and provided here.
power	Power parameter in the tree depth penalizing prior.
base	Base parameter in the tree depth penalizing prior.
fmu	Prior parameter for the center of the mean model.
k	Prior parameter for the mean model.
tau	Desired SD/nrtree for f function leaf prior if known.
dist	Distribution to be passed to intercept-only AFT model to center $y.train$ .
total.lambda	A rudimentary estimate of the process standard deviation. Used in calibrating the variance prior.
total.nu	Shape parameter for the variance prior.
mask	If a proportion is provided, then said quantile of $\max_i sd(x_i)$ is used to mask non-stationary departures (with respect to convergence) above this threshold.
K	Number of grid points for which to estimate survival probability.
events	Grid points for which to estimate survival probability.
TSVS	Setting to TRUE will avoid unnecessary processing for Thompson sampling variable selection, i.e., all that is needed is the variable counts from the tree branch decision rules.
drawDPM	Whether to utilize DPM or not.
alpha	Initial value of DPM concentration parameter.
alpha.a	Gamma prior parameter setting for DPM concentration parameter where $E[\alpha]=\alpha.a/\alpha.b$ .
alpha.b	See alpha.a above.
alpha.draw	Whether to draw alpha or it is fixed at the initial value.
neal.m	The number of additional atoms for Neal 2000 DPM algorithm 8.
constrain	Whether to perform constrained DPM or unconstrained.
m0	Center of the error distribution: defaults to zero.
k0.a	First Gamma prior argument for $k_0$ .
k0.b	Second Gamma prior argument for $k_0$ .
k0	Initial value of $k_0$ .
k0.draw	Whether to fix $k_0$ or draw it if from the DPM LIO prior hierarchy: $k_0 \sim \text{Gamma}(k_0.a, k_0.b)$ , i.e., $E[k_0]=k_0.a/k_0.b$ .
a0	First Gamma prior argument for $\tau$ .
b0.a	First Gamma prior argument for $b_0$ .
b0.b	Second Gamma prior argument for $b_0$ .

<code>b0</code>	Initial value of <code>b0</code> .
<code>b0.draw</code>	Whether to fix <code>b0</code> or draw it from the DPM LIO prior hierarchy: $b0 \sim \text{Gamma}(b0.a, b0.b)$ , i.e., $E[b0] = b0.a/b0.b$ .
<code>na.rm</code>	Value to be passed to the <code>predict</code> function.
<code>probs</code>	Value to be passed to the <code>predict</code> function.
<code>printevery</code>	Outputs MCMC algorithm status every <code>printevery</code> iterations.
<code>transposed</code>	Specify TRUE if all of the pre-processing for <code>xftrain/xstrain/xfctest/xstest</code> has been conducted prior to the call (including transposing).
<code>pred</code>	Specify TRUE if you want to return the <code>pred</code> item that is used to calculate <code>soffset</code> .

### Details

`nft2()/nft()` is the function to fit time-to-event data. The most general form of the model allowed is  $Y(\mathbf{x}) = \mu + f(\mathbf{x}) + sd(\mathbf{x})Z$  where  $E$  follows a nonparametric error distribution by default.

The `nft2()/nft()` function returns a fit object of S3 class type `nft2/nft` that is essentially a list containing the following items.

### Value

<code>ots,oid,ovar,oc,otheta</code>	These are XPtrs to the BART $f(x)$ objects in RAM that are only available for fits generated in the current R session.
<code>sts,sid,svar,sc,stheta</code>	Similarly, these are XPtrs to the HBART $sd(x)$ objects.
<code>fmu</code>	The constant $\mu$ .
<code>f.train,s.train</code>	The trained $f(x)$ and $sd(x)$ respectively: matrices with <code>ndpost</code> rows and $n$ columns.
<code>f.train.mean,s.train.mean</code>	The posterior mean of the trained $f(x)$ and $sd(x)$ respectively: vectors of length $n$ .
<code>f.trees,s.trees</code>	Character strings representing the trained fits of $f(x)$ and $sd(x)$ respectively to facilitate usage of the <code>predict</code> function when XPtrs are unavailable.
<code>dpalph</code>	The draws of the DPM concentration parameter $\alpha$ .
<code>dpn,dpn.</code>	The number of atom clusters per DPM, $J$ , for all draws including burn-in and excluding burn-in respectively.
<code>dpmu</code>	The draws of the DPM parameter $\mu[i]$ where $i = 1, \dots, n$ indexes subjects: a matrix with <code>ndpost</code> rows and $n$ columns.
<code>dpmu.</code>	The draws of the DPM parameter $\mu[j]$ where $j = 1, \dots, J$ indexes atom clusters: a matrix with <code>ndpost</code> rows and $J$ columns.
<code>dpwt.</code>	The weights for efficient DPM calculations by atom clusters (as opposed to subjects) for use with <code>dpmu.</code> (and <code>dpsd.</code> ; see below): a matrix with <code>ndpost</code> rows and $J$ columns.

dpsd, dpsd.	Similarly, the draws of the DPM parameter $\tau[i]$ transformed into the standard deviation $\sigma[i]$ for convenience.
dpC	The indices $j$ for each subject $i$ corresponding to their shared atom cluster.
z.train	The data values/augmentation draws of $\log t$ .
f.tmind/f.tavgd/f.tmaxd	The min/average/max tier degree of trees in the $f$ ensemble.
s.tmind/s.tavgd/s.tmaxd	The min/average/max tier degree of trees in the $s$ ensemble.
f.varcount, s.varcount	Variable importance counts of branch decision rules for each $x$ of $f$ and $s$ respectively: matrices with <code>ndpost</code> rows and $p$ columns.
f.varcount.mean, s.varcount.mean	Similarly, the posterior mean of the variable importance counts for each $x$ of $f$ and $s$ respectively: vectors of length $p$ .
f.varprob, s.varprob	Similarly, re-weighting the posterior mean of the variable importance counts as sum-to-one probabilities for each $x$ of $f$ and $s$ respectively: vectors of length $p$ .
LPML	The log Pseudo-Marginal Likelihood as typically calculated for right-/left-censoring.
pred	The object returned from the <code>predict</code> function where <code>x.test=x.train</code> in order to calculate the <code>soffset</code> item that is needed to use <code>predict</code> when <code>XPtrs</code> are not available.
soffset	See <code>pred</code> above.
aft	The AFT model fit used to initialize NFT BART.
elapsed	The elapsed time of the run in seconds.

### Author(s)

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### References

Sparapani R., Logan B., Maiers M., Laud P., McCulloch R. (2023) Nonparametric Failure Time: Time-to-event Machine Learning with Heteroskedastic Bayesian Additive Regression Trees and Low Information Omnibus Dirichlet Process Mixtures *Biometrics (ahead of print)* <doi:10.1111/biom.13857>.

### See Also

[predict.nft2](#), [predict.nft](#)

### Examples

```
##library(nftbart)
data(lung)
N=length(lung$status)

##lung$status: 1=censored, 2=dead
```

```

##delta: 0=censored, 1=dead
delta=lung$status-1

## this study reports time in days rather than weeks or months
times=lung$time
times=times/7 ## weeks

## matrix of covariates
x.train=cbind(lung[, -(1:3)])
## lung$sex:      Male=1 Female=2

## token run just to test installation
post=nft2(x.train, x.train, times, delta, K=0,
          nskip=0, ndpost=10, nadapt=4, adaptevery=1)

set.seed(99)
post=nft2(x.train, x.train, times, delta, K=0)
XPtr=TRUE

x.test = rbind(x.train, x.train)
x.test[, 2]=rep(1:2, each=N)
K=75
events=seq(0, 150, length.out=K+1)
pred = predict(post, x.test, x.test, K=K, events=events[-1],
              XPtr=XPtr, FPD=TRUE)

plot(events, c(1, pred$surv.fpd.mean[1:K]), type='l', col=4,
      ylim=0:1,
      xlab=expression(italic(t)), sub='weeks',
      ylab=expression(italic(S)(italic(t), italic(x))))
lines(events, c(1, pred$surv.fpd.upper[1:K]), lty=2, lwd=2, col=4)
lines(events, c(1, pred$surv.fpd.lower[1:K]), lty=2, lwd=2, col=4)
lines(events, c(1, pred$surv.fpd.mean[K+1:K]), lwd=2, col=2)
lines(events, c(1, pred$surv.fpd.upper[K+1:K]), lty=2, lwd=2, col=2)
lines(events, c(1, pred$surv.fpd.lower[K+1:K]), lty=2, lwd=2, col=2)
legend('topright', c('Adv. lung cancer\nmortality example',
                    'M', 'F'), lwd=2, col=c(0, 4, 2), lty=1)

```

---

predict.aftree

*Estimating the survival and the hazard for AFT BART models.*


---

## Description

The function `predict.aftree()` is provided for performing posterior inference via test data set estimates stored in a `aftree` object returned from `AFTree()` in a similar fashion as that of `predict.nft`. N.B. the `x.test` matrix must be provided on the `AFTree()` function call. Here we are only calculating the survival function by default, and, if requested, the hazard as well.

**Usage**

```
## S3 method for class 'aftree'
predict(
  ## data
  object,
  ## predictions
  events=NULL,
  FPD=FALSE,
  probs=c(0.025, 0.975),
  take.logs=TRUE,
  seed=NULL,
  ## default settings
  ndpost=nrow(object$mix.prop),
  nclust=ncol(object$mix.prop),
  ## etc.
  ...)
```

**Arguments**

object	Object of type <code>nft</code> from a previous call to <code>nft()</code> .
events	You must specify a grid of time-points; however, they can be a matrix with rows for each subject.
FPD	Whether to yield the usual predictions or marginal predictions calculated by the partial dependence function.
probs	A vector of length two containing the lower and upper quantiles to be calculated for the predictions.
take.logs	Whether or not to take logarithms.
seed	If provided, then this value is used to generate random natural logarithms of event times from the predictive distribution.
ndpost	The number of MCMC samples generated.
nclust	The number of DPM clusters generated.
...	The et cetera objects passed to the <code>predict</code> method. Currently, it has no functionality.

**Details**

Returns a list with the following entries. If `hazard=TRUE` is specified, then a similar set of entries for the hazard are produced.

**Value**

surv.fpd	Survival function posterior draws on a grid of time-points by the partial dependence function when requested.
surv.fpd.mean	Survival function estimates on a grid of time-points by the partial dependence function when requested.



- surv.fpd.lower Survival function lower quantiles on a grid of time-points by the partial dependence function when requested.
- surv.fpd.upper Survival function upper quantiles on a grid of time-points by the partial dependence function when requested.

**Author(s)**

Rodney Sparapani: <rsparapa@mcw.edu>

**See Also**

[predict.nft](#)

---

predict.nft2

*Drawing Posterior Predictive Realizations for NFT BART models.*

---

**Description**

The function `predict.nft2()/predict.nft()` is the main function for drawing posterior predictive realizations at new inputs using a fitted model stored in a `nft2/nft` object returned from `nft2()/nft()`.

**Usage**

```
## S3 method for class 'nft2'
predict(
  ## data
  object,
  xftest=object$xftrain,
  xstest=object$xstrain,
  ## multi-threading
  tc=getOption("mc.cores", 1), ##OpenMP thread count
  ## current process fit vs. previous process fit
  XPtr=TRUE,
  ## predictions
  K=0,
  events=object$events,
  FPD=FALSE,
  probs=c(0.025, 0.975),
  take.logs=TRUE,
  na.rm=FALSE,
  RMST.max=NULL,
  ## default settings for NFT:BART/HBART/DPM
  fmu=object$NFT$fmu,
  soffset=object$soffset,
  drawDPM=object$drawDPM,
  ## etc.
```

```

...))

## S3 method for class 'nft'
predict(
  ## data
  object,
  x.test=object$x.train,
  ## multi-threading
  tc=getOption("mc.cores", 1), ##OpenMP thread count
  ## current process fit vs. previous process fit
  XPtr=TRUE,
  ## predictions
  K=0,
  events=object$events,
  FPD=FALSE,
  probs=c(0.025, 0.975),
  take.logs=TRUE,
  na.rm=FALSE,
  RMST.max=NULL,
  ## default settings for NFT:BART/HBART/DPM
  fmu=object$NFT$fmu,
  soffset=object$soffset,
  drawDPM=object$drawDPM,
  ## etc.
  ...)

```

### Arguments

<code>object</code>	Object of type <code>nft2/nft</code> from a previous call to <code>nft2()/nft()</code> .
<code>xfstest, xstest, x.test</code>	New input settings in the form of a matrix at which to construct predictions. Defaults to the training inputs.
<code>tc</code>	Number of OpenMP threads to use for parallel computing.
<code>XPtr</code>	If <code>object</code> was created during the currently running R process, then (via an Rcpp <code>XPtr</code> ) the BART/HBART tree ensemble objects can be accessed in RAM; otherwise, those objects will need to be loaded from their string encodings.
<code>K</code>	The length of the grid of time-points to be used for survival predictions. Set to zero to avoid these calculations which can be time-consuming for large data sets.
<code>events</code>	You can specify the grid of time-points; otherwise, they are derived from quantiles of the augmented event times.
<code>FPD</code>	Whether to yield the usual predictions or marginal predictions calculated by the partial dependence function.
<code>probs</code>	A vector of length two containing the lower and upper quantiles to be calculated for the predictions.
<code>take.logs</code>	Whether or not to take logarithms.

na.rm	Whether NA values should be removed from the summaries.
RMST.max	To calculate Restricted Mean Survival Time (RMST), we need to set a reasonable time maxima. Typically, a clinically important time that a majority (or a large plurality) of censored subjects have been followed through that point or beyond.
fmu	BART centering parameter for the test data. Defaults to the value used by nft2()/nft() when training the model.
soffset	HBART centering parameter for the test data. Defaults to the value used by nft2()/nft() when training the model.
drawDPM	Whether NFT BART was fit with, or without, DPM.
...	The et cetera objects passed to the predict method. Currently, it has no functionality.

### Details

predict.nft2()/predict.nft() is the main function for calculating posterior predictions and uncertainties once a model has been fit by nft2()/nft().

Returns a list with the following entries.

### Value

f.test	Posterior realizations of the mean function stored in a matrix. Omitted if partial dependence functions are performed since these will typically be large.
s.test	Posterior realizations of the SD function stored in a matrix. Omitted if partial dependence functions are performed since these will typically be large.
f.test.mean	Posterior predictive mean of mean function.
f.test.lower	Posterior predictive lower quantile of mean function.
f.test.upper	Posterior predictive upper quantile of mean function.
s.test.mean	Posterior predictive mean of SD function.
s.test.lower	Posterior predictive lower quantile of SD function.
s.test.upper	Posterior predictive upper quantile of SD function.
surv.fpd	Survival function posterior draws on a grid of time-points by the partial dependence function when requested.
surv.fpd.mean	Survival function estimates on a grid of time-points by the partial dependence function when requested.
surv.fpd.lower	Survival function lower quantiles on a grid of time-points by the partial dependence function when requested.
surv.fpd.upper	Survival function upper quantiles on a grid of time-points by the partial dependence function when requested.

### Author(s)

Rodney Sparapani: <rsparapa@mcw.edu>

**See Also**[nft2](#), [nft](#)

---

`tsvs2`*Variable selection with NFT BART models.*

---

**Description**

The `tsvs2()/tsvs()` function is for Thompson sampling variable selection with NFT BART.

**Usage**

```

tsvs2(
  ## data
  xftrain, xstrain, times, delta=NULL,
  rm.const=TRUE, rm.dupe=TRUE,
  ##tsvs args
  K=20, a.=1, b.=0.5, C=0.5,
  rds.file='tsvs2.rds', pdf.file='tsvs2.pdf',
  ## multi-threading
  tc=getOption("mc.cores", 1), ##OpenMP thread count
  ##MCMC
  nskip=1000, ndpost=2000,
  nadapt=1000, adaptevery=100,
  chvf=NULL, chvs=NULL,
  method="spearman", use="pairwise.complete.obs",
  pbd=c(0.7, 0.7), pb=c(0.5, 0.5),
  stepwpert=c(0.1, 0.1), probchv=c(0.1, 0.1),
  minnumbot=c(5, 5),
  ## BART and HBART prior parameters
  ntree=c(10, 2), numcut=100,
  xifcuts=NULL, xiscuts=NULL,
  power=c(2, 2), base=c(0.95, 0.95),
  ## f function
  fmu=NA, k=5, tau=NA, dist='weibull',
  ## s function
  total.lambda=NA, total.nu=10, mask=0.95,
  ## survival analysis
  ##K=100, events=NULL,
  ## DPM LIO
  drawDPM=1L,
  alpha=1, alpha.a=1, alpha.b=0.1, alpha.draw=1,
  neal.m=2, constrain=1,
  m0=0, k0.a=1.5, k0.b=7.5, k0=1, k0.draw=1,
  a0=3, b0.a=2, b0.b=1, b0=1, b0.draw=1,
  ## misc
  na.rm=FALSE, probs=c(0.025, 0.975), printevery=100,

```

```

    transposed=FALSE
  )
  tsvs(
    ## data
    x.train, times, delta=NULL,
    rm.const=TRUE, rm.dupe=TRUE,
    ##tsvs args
    K=20, a.=1, b.=0.5, C=0.5,
    rds.file='tsvs.rds', pdf.file='tsvs.pdf',
    ## multi-threading
    tc=getOption("mc.cores", 1), ##OpenMP thread count
    ##MCMC
    nskip=1000, ndpost=2000,
    nadapt=1000, adaptevery=100,
    chv=NULL,
    method="spearman", use="pairwise.complete.obs",
    pbd=c(0.7, 0.7), pb=c(0.5, 0.5),
    stepwpert=c(0.1, 0.1), probchv=c(0.1, 0.1),
    minnumbot=c(5, 5),
    ## BART and HBART prior parameters
    ntree=c(10, 2), numcut=100, xicuts=NULL,
    power=c(2, 2), base=c(0.95, 0.95),
    ## f function
    fmu=NA, k=5, tau=NA, dist='weibull',
    ## s function
    total.lambda=NA, total.nu=10, mask=0.95,
    ## survival analysis
    ##K=100, events=NULL,
    ## DPM LIO
    drawDPM=1L,
    alpha=1, alpha.a=1, alpha.b=0.1, alpha.draw=1,
    neal.m=2, constrain=1,
    m0=0, k0.a=1.5, k0.b=7.5, k0=1, k0.draw=1,
    a0=3, b0.a=2, b0.b=1, b0=1, b0.draw=1,
    ## misc
    na.rm=FALSE, probs=c(0.025, 0.975), printevery=100,
    transposed=FALSE
  )

```

### Arguments

xfttrain	n x pf matrix of predictor variables for the training data.
xstrain	n x ps matrix of predictor variables for the training data.
x.train	n x ps matrix of predictor variables for the training data.
times	nx1 vector of the observed times for the training data.

<code>delta</code>	nx1 vector of the time type for the training data: 0, for right-censoring; 1, for an event; and, 2, for left-censoring.
<code>rm.const</code>	To remove constant variables or not.
<code>rm.dupe</code>	To remove duplicate variables or not.
<code>K</code>	The number of Thompson sampling steps to take. Not to be confused with the size of the time grid for survival distribution estimation.
<code>a.</code>	The prior parameter for successes of a Beta distribution.
<code>b.</code>	The prior parameter for failures of a Beta distribution.
<code>C</code>	The probability cut-off for variable selection.
<code>rds.file</code>	File name to store RDS object containing Thompson sampling parameters.
<code>pdf.file</code>	File name to store PDF graphic of variables selected.
<code>tc</code>	Number of OpenMP threads to use.
<code>nskip</code>	Number of MCMC iterations to burn-in and discard.
<code>ndpost</code>	Number of MCMC iterations kept after burn-in.
<code>nadapt</code>	Number of MCMC iterations for adaptation prior to burn-in.
<code>adaptevery</code>	Adapt MCMC proposal distributions every adaptevery iteration.
<code>chvf, chvs, chv</code>	Predictor correlation matrix used as a pre-conditioner for MCMC change-of-variable proposals.
<code>method, use</code>	Correlation options for change-of-variable proposal pre-conditioner.
<code>pbd</code>	Probability of performing a birth/death proposal, otherwise perform a rotate proposal.
<code>pb</code>	Probability of performing a birth proposal given that we choose to perform a birth/death proposal.
<code>stepwpert</code>	Initial width of proposal distribution for perturbing cut-points.
<code>probchv</code>	Probability of performing a change-of-variable proposal. Otherwise, only do a perturb proposal.
<code>minnumbot</code>	Minimum number of observations required in leaf (terminal) nodes.
<code>ntree</code>	Vector of length two for the number of trees used for the mean model and the number of trees used for the variance model.
<code>numcut</code>	Number of cutpoints to use for each predictor variable.
<code>xifcuts, xiscuts, xicuts</code>	More detailed construction of cut-points can be specified by the <code>xicuts</code> function and provided here.
<code>power</code>	Power parameter in the tree depth penalizing prior.
<code>base</code>	Base parameter in the tree depth penalizing prior.
<code>fmu</code>	Prior parameter for the center of the mean model.
<code>k</code>	Prior parameter for the mean model.
<code>tau</code>	Desired SD/ <code>ntree</code> for f function leaf prior if known.
<code>dist</code>	Distribution to be passed to intercept-only AFT model to center <code>y.train</code> .

<code>total.lambda</code>	A rudimentary estimate of the process standard deviation. Used in calibrating the variance prior.
<code>total.nu</code>	Shape parameter for the variance prior.
<code>mask</code>	If a proportion is provided, then said quantile of $\max_i \text{sd}(x_i)$ is used to mask non-stationary departures (with respect to convergence) above this threshold.
<code>drawDPM</code>	Whether to utilize DPM or not.
<code>alpha</code>	Initial value of DPM concentration parameter.
<code>alpha.a</code>	Gamma prior parameter setting for DPM concentration parameter where $E[\alpha]=\text{alpha.a}/\text{alpha.b}$ .
<code>alpha.b</code>	See <code>alpha.a</code> above.
<code>alpha.draw</code>	Whether to draw <code>alpha</code> or it is fixed at the initial value.
<code>neal.m</code>	The number of additional atoms for Neal 2000 DPM algorithm 8.
<code>constrain</code>	Whether to perform constrained DPM or unconstrained.
<code>m0</code>	Center of the error distribution: defaults to zero.
<code>k0.a</code>	First Gamma prior argument for <code>k0</code> .
<code>k0.b</code>	Second Gamma prior argument for <code>k0</code> .
<code>k0</code>	Initial value of <code>k0</code> .
<code>k0.draw</code>	Whether to fix <code>k0</code> or draw it if from the DPM LIO prior hierarchy: $k0 \sim \text{Gamma}(k0.a, k0.b)$ , i.e., $E[k0]=k0.a/k0.b$ .
<code>a0</code>	First Gamma prior argument for <code>tau</code> .
<code>b0.a</code>	First Gamma prior argument for <code>b0</code> .
<code>b0.b</code>	Second Gamma prior argument for <code>b0</code> .
<code>b0</code>	Initial value of <code>b0</code> .
<code>b0.draw</code>	Whether to fix <code>b0</code> or draw it from the DPM LIO prior hierarchy: $b0 \sim \text{Gamma}(b0.a, b0.b)$ , i.e., $E[b0]=b0.a/b0.b$ .
<code>na.rm</code>	Value to be passed to the <code>predict</code> function.
<code>probs</code>	Value to be passed to the <code>predict</code> function.
<code>printevery</code>	Outputs MCMC algorithm status every <code>printevery</code> iterations.
<code>transposed</code>	<code>tsvs</code> handles all of the pre-processing for <code>x.train/x.test</code> (including transposing) computational efficiency.

## Details

`tsvs2()/tsvs()` is the function to perform variable selection.

The `tsvs2()/tsvs()` function returns a fit object of S3 class type `list` as well as storing it in `rds.file` for sampling in progress.

## Author(s)

Rodney Sparapani: <rsparapa@mcw.edu>

## References

- Sparapani R., Logan B., Maiers M., Laud P., McCulloch R. (2023) Nonparametric Failure Time: Time-to-event Machine Learning with Heteroskedastic Bayesian Additive Regression Trees and Low Information Omnibus Dirichlet Process Mixtures *Biometrics (ahead of print)* <doi:10.1111/biom.13857>.
- Liu Y., Rockova V. (2021) Variable selection via Thompson sampling. *Journal of the American Statistical Association. Jun 29:1-8.*

## See Also

[tsvs](#)

## Examples

```
##library(nftbart)
data(lung)
N=length(lung$status)

##lung$status: 1=censored, 2=dead
##delta: 0=censored, 1=dead
delta=lung$status-1

## this study reports time in days rather than weeks or months
times=lung$time
times=times/7 ## weeks

## matrix of covariates
x.train=cbind(lung[, -(1:3)])
## lung$sex:      Male=1 Female=2

##vars=tsvs2(x.train, x.train, times, delta)
vars=tsvs2(x.train, x.train, times, delta, K=0) ## K=0 just returns 0
```

---

xicuts

*Specifying cut-points for the covariates*

---

## Description

This function allows you to create a list that specifies the cut-points for the covariates.

## Usage

```
xicuts(x.train, transposed=FALSE, numcut=100)
```



**Arguments**

<code>x.train</code>	The training matrix to derive cut-points from.
<code>transposed</code>	Whether or not the matrix has been tranposed yet.
<code>numcut</code>	The number of cut-points to create.

**Details**

The cut-points are generated uniformly from `min.` to `max.`, i.e., the distribution of the data is ignored.

**Value**

An object is returned of type `BARTcutinfo` which is essentially a list.

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